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(54) Tide: STEROID MODIFIED OLIGONUCLEOTIDES

(57) Abstract

Oligonucleotides modified at their backbones by the attachment of at least one steroid are described. The modified oligon-ucleotides anchor in the cell membrane to serve as a probe and to provide therapeutic activity.

* See back of page

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STEROID MODIFIED OLIGONUCLEOTIDES

GRANT REFERENCE

5 This invention was developed with support provided by the National Cooperative Drug Discovery Group for the Treatment of AIDS, Grant U01 A124846 from the National Cancer Institute of Allergy and Infectious Disease and by Grant 5R37GM10265 from the National Institute of General Medical Science.

FIELD OF INVENTION, BACKGROUND AND PRIOR ART

This invention relates to oligonucleotides

15 modified by a pendant steroid group. More

particularly, the present invention is related to

steroid modified oligonucleotides and a method of

using the modified oligonucleotides as antiviral

agents.

- The pioneering work of Zamecnik and Stephenson, <u>Proc. Natl. Acad.</u>, <u>75</u>:280-284 (1978), on antiviral activity of oligonucleotides and Miller and Ts'o, on the chemistry and biochemistry of non-ionic analogues (Barrett, et al., <u>Biochem.</u>, <u>13</u>:4898-5
- 25 (1974) and Jayaraman, et al. <u>Proc. Natl. Acad. Sci.</u>
 <u>USA</u>, 78:1537-1541 (1981)) has stimulated extensive

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research directed at the therapeutic potential of nucleotide polymers. Oligonucleotide analogues with methylphosphonate, Miller, et al., <u>Biochemie</u>, <u>67</u>:769-776 (1985), Agris, et al., Biochem., <u>25</u>:6268-6275

- 5 (1986), Smith et al., Proc. Natl. Acad. Sci. USA, 83:2787-2791 (1986), and Sarin, et al., Proc. Natl. Acad. Sci. USA, 85:7448-7451 (1988); phosphorothicate, Matsukura, et al., Proc. Natl. Acad. Sci. USA, 85:7079-7083 (1988); and
- phosphoramidate, Agrawal, et al., Proc. Natl. Acad.

 Sci. USA, 85:7079-7083 (1988), backbones as well as natural type oligonucleotides, Zamecnik, et al.,

 Proc. Natl. Acad. Sci. USA, 83:4143-4146 (1986), and a polylysine conjugate, Goodchild, et al., Proc.
- Natl. Acad. Sci. USA, 85:5507-5511 (1988), have now been reported to inhibit viral replication in cell culture. The viruses studied in this context include Rous sarcoma virus, Samecnik and Stephenson, Proc.

 Natl. Acad., 75:280-284 (1978); simian virus, Miller,
- et al., Biochemie, 67: 769-776 (1985); vesticular stomatitis virus, Agris, et al., Biochem., 25:6268-6275 (9186) and Lemaitre, et al., Proc. Natl. Acad. Sci. USA, 84:648-652 (1987); human immunodeficiency virus (HIV), Sarin, et al., Proc. Natl. Acad. Sci.
- 25 <u>USA</u>, <u>85</u>:7448-7451 (1988), Matsukura, et al., <u>Proc.</u>
 Natl. Acad. Sci. USA, <u>84</u>:7706-7710 (1987), Agrawal,

et al., <u>Proc. Natl. Acad. Sci. USA</u>, <u>85</u>:7079-7083
(1988), Aamecnik, et al., <u>Proc. Natl. Acad. Sci. USA</u>,
<u>83</u>:4143-4146 (1986), and Goodchild, et al., <u>Proc. Natl. Acad. Sci. USA</u>,
<u>85</u>:5507-5511 (1988); herpes

5 simplex virus, Smith, et al., <u>Proc. Natl. Acad. Sci. USA</u>, <u>83</u>:2787-2791 (1986); and influenza virus,
Zerial, et al., <u>Nuc. Acids. Res.</u>, <u>15</u>:9909-9919
(1987).

The concept underlying this work is that an 10 oligonucleotide complementary to a unique segment of a viral genome, or an RNA derived from it, may selectively disrupt processes dependent on that segment by hybridization. This rationale is supported by a variety of experiments with cell free 15 systems or with cells to which "antisense" polynucleotides have been inserted by microinjection or transfection, C.A. & Cohen, J.S., Cancer Res., 48:2659-2668 (1988). However, the actual mechanisms by which oligonucleotides and their analogs function 20 as inhibitors in cell cultures are still far from clear. In particular, little is known about the interaction of the oligomers with cell membranes or the locus of their reactions within cells. It appears that non-ionic oligomers, such as the methyl 25 phosphonate analogues diffuse passively through cell membranes.

S.E. Clare has synthesized oligonucleotides possessing one or more 2,2,2,-trichloral-1,1-dimethylethyl (TDCME, lipophilic) group of the phosphorous atom in the chain and show that this group on one strand with proper stereochemistry can inhibit cleavage of the opposite strand by a restriction endonuclease and that the same group on a template will inhibit synthesis of the complementary strand by the Klenow enzyme. S.E. Clare also demonstrated a single modification 5' to dGNAd (CG) octamer by TDCME group prevents the B to Z conformational transition. S.E. Clare, Ph.D. Dissertation, Northwestern University, Evanston, Illinois (1987).

family of oligonucleotides modified at the backbone so that the oligonucleotide may anchor at the cell membrane to provide antiviral effects. The present invention describes a family of oligonucleotides with a modification designed to anchor the oligomer, at least transiently at the cell membrane, to inhibit HIV-1 in cell culture. Fatty substances have been selected as an anchor for the oligonucleotide, and without being limitative, steroids such as

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anchor since they are highly hydrophobic and cell membranes have an abundance of this steriod. The compounds may also have anti-sense activity.

The cholesteryl is a large lipophilic 5 group, much larger than the TDCME group. principle, such pendent groups, when linked covalently to the internucleotide phosphorous atoms, have potential as lipophilic centers to enhance the interaction with membranes, to alter partitioning of 10 oligonucleotides within cells, to inhibit certain enzymatic reactions and to influence the stability of hybrids joined with natural polynucleotides. Cholesteryl is a component of any biological membrane and interacts with other lipids. The AIDS virus, 15 HIV, is distinguished by an unusually high cholesteryl content in the lipid membrane. Early model studies by Finean, Experientia, 9:17-19 (1985), suggested that the cholesteryl molecule is capable of formation of a stabilizing complex with the

phospholipid molecule. The hydrocarbon chain of the cholesteryl is bound to the parallel portion of the phospholipid chain by Van DeWall forces. Recent studies employing a variety of techniques indicated that the major forces may involve the hydrophobic

portion of the lipid molecules. Therefore, cholesteryl is a preferred modifying group for oligonucleotide interaction with cells.

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SUMMARY OF THE INVENTION

The invention is concerned with oligonucleotides that are modified to anchor the oligomer at the cell membrane or eventually,

intranuclearly, so that the oligonucleotide may serve as a probe and provide therapeutic activity.

More specifically, the invention provides a pharmaceutical composition including an oligonucleotide conjugated to a steroid for increasing the antiviral activity of the compound. One aspect of the present invention provides oligonucleotide compounds represented by the following structural formula:

Formula I

O R O
O R O
O P N - A - N - C - O - Cholesteryl
O R
NUC

10

5

wherein A = aliphatic alkyl or branched aliphatic alkyl or a heteroatom containing an alkyl (branched) chain of 2 to 18 carbon atoms, preferably CH₂, R = H and lower alkyl up to 12 carbon atoms, preferably methyl; NUC refers to an oligonucleotide which may be a deoxyribonucleotide or a ribonucleosite. Preferred nucleotides are ethymidine, deoxyadenosine, deoxyguanosine and deoxycytidine. The nucleotides are connected respectively to the phosphorous through their 3' and 5' oxygens, B is a purine or pyrimidine base (such as Thy, Cyt, Gua, Ade).

Cholesteryl has been selected as the preferred anchol because it is highly hydrophilic and found in cell membranes. The cholesteryl modified oligonucleotides of the present invention have been

found to inhibit HIV-1 in cell culture. The location of the insertion of the cholesteryl anchor on the oligonucleotide may be varied and is not dependent on sequence. However, other steriods are efficacious

5

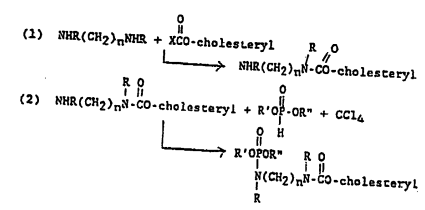
DETAILED DESCRIPTION OF THE INVENTION

The novel compounds of Formula I can be prepared by convenient procedures for introducing a cholesteryl group at any desired internucleoside phosphorous in the course of synthesizing an oligonucleotide. The cholesteryl may be linked to an oligonucleotide as a substituent at either the 3'-0 or 5'-0 terminus.

Processes for preparing the novel compounds of Formula I are generally described by equations A and B.

Δ.

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In the reactions, X = C1- and pnitrophenoxy, n = 2 and 6, and R = H and methyl. The
reactions for linking the amines to phosphorous are
based on the general procedure of Froehler, B.C.,
Tet. Lett., 27:5575-5578 (1986) for generating P-N
bonds in oligonucleotide derivatives. The article is
incorporated by reference.

Equation A avoids side reactions involving condensation at both nitrogen atoms of the diamine to form bis-phosphoramidates. Procedures for preparing fifteen compounds of Table I with the cholesteryl anchor are described as follows: phosphodiester links were formed by cyanoethyl phosphoramidite chemistry described in the standard synthesis protocol provided

by the manufacturer of the synthesizer, for example, Biosearch 8600, Biosearch, Inc., San Raphael, California. Chain extension by hydrogen phosphonate chemistry is described by Froehler, et al., Tet.

5 Lett., 27:469-472 (1986) and Froehler, et al., Nuc. Acids. Res., 14:5399-5407 (1986). Phosphorothioate functional groups are added by the procedure of Froehler, et al., Tet. Lett., 27:5575-5578 (1986).

10 Experimental Procedure

2-(Cholesteryloxycarbonylamino) ethylamine.

Cholesteryl chloroformate (2g) in dichloromethane (6 ml) was added dropwise to a solution of ethylenediamine (2.5 ml) in dichloromethane (6 ml) and pyridine (6 ml). The mixture was stirred for two hours; then the solvent was removed under vacuum and the residue was partitioned between water (150 ml) and dichloromethane (150 ml). The organic layer was washed with water, dried (Na₂SO₄), and concentrated to give the title compound; 1.6 g (76%), mp 149-155 degrees Centigrade. Recrystallization from cyclohexane afforded crystales melting at 152-155 degrees Centigrade; Rf on silica (CHC₁₃/MeOH, 1/1

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v/v) 0.15; positive ninhydrin test. Anal. Calcd for $^{\rm C}_{30}{}^{\rm H}_{52}{}^{\rm N}_{2}{}^{\rm O}_{2}$; C, 76.22; H, 11.09; N, 5.93. Found: c, 75.96; H, 11.10; N, 5.86.

Preparation of Cholesteryl-Modified Dinucleoside Monophosphate on CPG Support.

Internucleoside cholesteryl side chains were linked to phosphorous via phosphoramidate bonds (adaption of procedure of Froehler). The preparation 10 of d-DMT-ibGcibG-CPG. is representative. A sample of DMT-ibG linked through the 3'-0 to a controlled-porglass support (Biosearch) (250 mg. 8 micro moles of DMG-ibG) was placed in a Glencoe Gastight syringe (10 ml) equipped with a plug of glass wool at the inlet. 15 Reactions and washings were carried out by drawing in and ejecting the desired solutions. Thus, the DMT groups were removed with dichloroacetic acid (2.5% in CH2Cl2; the support was washed repeatedly with $C_{\eta}H_{\eta}/CH_{\eta}CN$ (1/4), coupling was effected by drawing in 20 together DMT-ibG-hydrogen phosphonate (80 mg., 0.1 mmol, in 1.2 ml CH_3CN/C_5H_5) and trimethylacetyl chloride (65 micro L, 0.5 micro moles, in 1.2 ml CH_3CN/C_5H_5 , 1/1 v/v; 2 minutes), and the support was washed well with CH_3CN/C_5H_5 . A solution of

cholesteryloxycarbonylaminoethylamine (250 mg, 0.5 mmol) in CCl $_4$ (5 ml) and C $_5\mathrm{H}_5$ (2 ml) was then drawn

into the syringe and after 0.5 hours, the solution was ejected and the solid was washed well with CH₃CN. Appropriate portions were then transferred to a cartridge for extension by machine synthesis (Biosearch 8600 Synthesizer) or to a syringe for manual synthesis.

Chain Extension. The oligonucleotide chains were extended by conventional phosphoramidite chemistry in constructing phosphodiester links and by hydrogen phosphonate chemistry in building the phosphorothicate derivatives. The manual procedure used in adding a thymidine unit to DMT-ibG*G-CPG in synthesizing compound 2 in Table 1 is representative of one synthetic cycle utilizing a phosphoramidite reagent.

The DMT(G*G) loaded CPG (30 mg, 1 micro mole) was poured into a 1.0 ml Glenco Gas tight syringe with a glass wool plug at the inlet. Washes were effected by drawing up the desired amount of the reagent, resuspending the support by brief hand agitation, and ejecting the solution. The DMT protecting group was removed by washing with DCA/CH₂Cl₂ (2.5/100 v/v, 5.0 ml), and organ effluents were pooled from subsequent spectroscopy (447 nm) and calculation of the coupling efficiency. The support was washed successively with C₅H₅N/CH₃CN (1/4, v/v, 1

X 0.5 ml), and CH₃CN (2 X 0.5 ml). Any unreacted 5'-OH groups were capped by drawing DMAP in C₅H₅N/THF (0.3 M, 1/15, v.v, 0.5 ml) into the syringe followed immediately by Ac₂O/THF (0.6 M, 0.5 ml). The mixture was agitated for one minute, capping agents were ejected from the syringe, and the support was washed with C₅H₅N/CH₃CN (1/4, v/v, 1 X 5.0 ml) and CH₃CN (1X 0.5 ml). The phosphite internucleoside linkage was oxidized to the phosphotriester linkage with I₂ in C₅H₅N/THF H₂O (0.1 M I₂, 18/80/2, v/v/v, 0.5 ml) for two minutes. The oxidant was ejected, and the support was washed with C₅H₅N/CH₃CN (1/4, v/v, 3 X 1.5 ml) and CH₃CN (3 X 1.5 ml) to complete one synthetic cycle.

chemistry the DMT-ibG*ibG-cpg (1 micro mole loaded dimer) was detritylate as in the previous case. A solution of the DMT-nucleoside hydrogen phosphonate (10 mg, about 15 micro moles, in CH₃CN/C₅H₅N (1/1, v/v, 0.3 ml) was drawn into the syringe, which was agitated for two minutes. The coupling agents were ejected from the syringe, and the support washed with C₅H₅N/CH₃CN (1/1, v/v, 0.5 ml), and CH₃CN (3 X 0.5 ml) to complete one synthetic cycle. Additional couplings were performed by returning to the initial wash and repeating the cycle. Oxidation following

the final coupling step was performed by treatment with 0.1 M sulfur in CCl_4/Et_3N (9/1, v/v) at room temperature (two hours reaction). Procedures for machine syntheses were similar.

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Isolation of Oligonucleotides

The oligomers were removed from the syringe or the synthesizer and warmed in a capped vessel with concentrated NH₄OH at 55 degrees Centigrade for five hours. The aqueous solution was then removed and concentrated under reduced pressure to give the crude oligonucleotide. This substance was chromatographed on a C-18 column and the band corresponding to the desired target oligomer was collected and

15 lyophilized.

Table 1. Properties of Oligonucleotides

COM	pound	HPLC ^a min	TLC ^b Rf	PAGEC R _{IB}	oC Tmd
1	ACACCCAATTUTGAAAATGC	12.2	0.26	0.64	60
	Cholesteryl Substituents				
2	Acacccaattctgaaaatc*c	46.0	0.41	0.55	60
3	Ac <u>i</u> cc <u>c</u> aa <u>a</u> ga <u>t</u> aa <u>a</u> g*c	46.6	0.43	0.54	
4	A*CACCCAATTCTGAAAATG*C	61.0	0.58	•	52
5	Caattctcaaaatg*g	46.5	0.54	0.64	46.
6	$\mathtt{O}_{\mathbf{g}}\mathtt{O}_{\mathbf{g}}\mathtt{A}_{\mathbf{g}}\mathtt{A}_{\mathbf{g}}\mathtt{A}_{\mathbf{g}}\mathtt{A}_{\mathbf{g}}\mathtt{T}_{\mathbf{g}}\mathtt{T}_{\mathbf{g}}\mathtt{T}_{\mathbf{g}}\mathtt{T}_{\mathbf{g}}\mathtt{A}_{\mathbf{g}}\mathtt{A}_{\mathbf{g}}\mathtt{A}_{\mathbf{g}}\mathtt{A}_{\mathbf{g}}\mathtt{A}_{\mathbf{g}}\mathtt{T}_{\mathbf{g}}$	16.5	. 0.59	0.67	44
7	$A_3C_5A_6C_5C_3C_3A_5A_5T_5T_5C_5T_5G_5A_5A_5A_5A_5T_5G^\pm G$	40.2	0.61	0.64	47.
8	$C_{S}A_{\mathtt{S}}A_{\mathtt{S}}T_{\mathtt{S}}T_{\mathtt{S}}C_{\mathtt{S}}T_{\mathtt{S}}G_{\mathtt{S}}A_{\mathtt{S}}A_{\mathtt{S}}A_{\mathtt{S}}T_{\mathtt{S}}G_{\mathtt{S}}G$	17.3	0.47	0.67	29
9	C _S A _S A _S T _S T _S C _S T _S G _S A _S A _S A _S T _S G*C	47.4	0.59	0.63	25
10	C _S T _S G _S A _S A _S A _S A _S T _S G*G	15.6	0.52	0.76	
11	C _S T _S G _S A _S A _S A _S A _S T _S G*G	49.6	0.56	0.62	
12	G _s A _s C _s T _s T _s T _s T _s A _s G*G	45	0.60	0.79	••
13	C _S T _S T _S T _S T _S T _S T _S O×C	45.2	0.59	0.76	••
14	А _Б А _Б А _Б А _Б С*С	47.5	0.66	U.62	••
15	IsIsIsIsIsIsIsIsIsIsIsIsIsIsI	50.0	0.60	0.65	16

In formulas * represents O=P-NH(CH₂)₂NHCO₂ Cholesteryl; s represents O=P-S⁻, and + represents O=PNH(CH₂)₃N(CH₃)₂. Altered nucleotides in 7 are underlined.

Elution time, Hewlett-Packard RP-C18 column (10 cm); 0.1 M triethylammonium acetate (pH 7.0), 1%/min acetonitrile gradient starting at 0% acetonitrile; 0.5 ml/min flow rate.

bThin layer chromatography on Merck silica 10 plates with propanol/ammonium hydroxide/water, 55/10/35 v/v/v.

 $^{\text{C}}\text{Polyacrylamide}$ gel electrophoresis in 20% crosslinked gel at pH 8.0, $R_{\underline{m}}$ is migration relative to bromophenol blue.

the maximum slope in a plot of A₂₆₀ versus temperature in 0.1 M aqueous NaCl, 0.01 M Tris buffer at pH 7.0; total nucleotide concentration (base units) in approximately 10⁻⁴ M. In each case, the complement is a phosphodiester strand equal in length to the modified oligomer.

 $^{\rm e}{\rm The}$ sample appeared as a broad streak starting at R $_{\rm m}$ 0.2. The complement for determination of T $_{\rm m}$ was poly d(A).

25

Coupling efficiency introducing a cholesteryl fragment to the compounds of Table 1 exceeded 50%.

As noted, phosphodiester links were formed

in compounds 2-5 by conventional cyanoethyl

phosphoramidaite chemistry as described in the

standard synthesis protocol provided by the

manufacturer of the synthesizer, Biosearch, Inc., San

Raphael, California. For compounds 6 - 15, the

chains were extended by hydrogen phosphonate

chemistry as described by Froehelr, et al., Tet.

Lett., 27:469-472 (1986) and Froehler, et al., Nuc.

Acids. Res., 14:5399-5407 (1986) the final oxidation

with sulfur to generate the phosphorothicate groups.

- The compounds were characterized by HPLC,
 TLC, PAGE, thermal disassociation curves for hybrids
 formed with complimentary strands and by UV and NMR
 spectroscopy. The NMR spectra exhibited the
 characteristic peaks for phosphodiester,
- phosphoramidate and phosphorothicate functional groups. A proton NMR spectrum of compound 2 shows the presence of the cholesteryl fragment. Further, the hydrophobic nature of the cholesteryl-oligonucleotides was shown by the HPLC data in Table
- 25 1. For example, the elution times for samples analyzed by a reverse phase C-18 column increased

from 14 to 46 to 61 minutes for the series 1 (control), 2(one) cholesteryl, and 4(two) cholesteryl, respectively. Susceptibility to nuclease degradation was examined with compound 2.

In the presence of snake venom, phosphodiesterase, an alkaline phosphatase, compound 2 was completely hydrolyzed to the expected nucleosides and the fragment corresponding to the terminal G*G.

The data in Table 1 shows that the introduction of a single cholesteryl fragment at a terminal internucleoside position has only a minor effect on the stability of the hybrid duplex as measured by $T_{\rm m}$ values when compound 1 is compared with 2; compound 6 compared with 7, and compound 8

15 compared with 9. Conversely, two cholesteryl substitutents led to appreciable destabilization for the 20-mer compound, compare compound 4 with 1. The disassociation of complexes formed from equi-molar quantities in modified and unmodified complementary

20 oligodeoxyribo-nucleotides were measured by changes in adsorbents in 260 nm as a function of temperature.

Additional compounds were made in accordance with Equation B, wherein $R = CH_3$, N = 6, $X = -0C_6H_5MO_2$. The following compounds were prepared:

25 16, TTTTTTTTTT; 17, T#TTTTTTTT; 18, TTTT#TTTTT; and 19, CGCG#AATTCGCG, where # is O = P-N

 $(CH_3)(CH_2)CO_2$ cholesteryl. The procedures of Froehler, B.C., <u>Tet. Lett.</u>, <u>27</u>:5575-5578 (1986) and Marcus=Sekura, et al., Nuc. Acids. Res., 15:5749-5763 (1987) were followed in preparing phosphodiester links in compounds with internal modifications to avoid or minimize complications which could arise from the premature formation of phosphodiester groups in the course of the synthesis. As in the case of the derivatives of ethylenediamine, introduction of substituents at the terminal internucleoside links had little effect on T values for the hybrids formed in complementary sequence (T_{m} for 1:1 complexes with poly d(A) in 0.1 M NaCl, pH 7: 28 degrees Centigrade for compound 16, 27 degrees Centigrade for compound 15 17 and 28 degrees Centigrade for parent T_qT). Conversely, modification of the centrally positioned internal link led to significant destabilization (21 degrees centigrade T_{m} for compound 18). The mobility of stereoisomers of compound 19 on HPCL differed 20 sufficiently to permit separation of the isomers. The T_m for the duplexes formed from these selfcomplementary modified strands (stereoisomers of compound 15) were substantially lower than that for the parent duplex (T_{m} :40 degrees and 45 degrees

25 Centigrade for the isomers as compared to 56 degrees

Centigrade for CGCGAATTCGCG; 0.1 M NaCl, pH 7.0).

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The linking of cholesteryl to an oligonucleotide as a substituent at the 5'-0 terminus is quite simple and can be shown by the following example.

5

Synthesis and characterization of cholestery1-sTsGsG.

DMT(G) loaded CPG (88.2 mg, about 3 micro moles) was poured into 5 ml Glencoe gas tight syringe with a glass wool plug at the inlet. Washes were effected by drawing up the desired amount of the reagent, resuspending the support by brief hand agitation and ejecting the solution. The support was initially washed with CH₃CN (4.5 ml X 3) and CH₂Cl₂ (4.5 ml X 2). The DMT protecting group was removed by washing with DCA/CH₂Cl₂ (2.5/100, v/v, 10 ml). All of the orange effluents were pooled for subsequent spectroscopy (448 nm) and calculation of the coupling efficiency.

The support was washed successively with

20 pyridine/CH₃CH (1/4, v/v, 4.5 ml X 3), CH₃CN (4.5 ml

X 4) and dry CH₃CN (4.5 ml X 6). The H-phosphonate
solution (for G and T - 36 mg, about 0.06 micro moles
in dry CH₃CH/pyridine, 1/1, v/v, 2ml); for
cholesteryl H-phosphonate-42 mg., about 0.09 micro

25 moles in dry CH₃CN/pyridine, 1/3, v/v, 2.0 ml) and
trimethylacetyl chloride solution (0.03 ml, about

0.351 micro moles in trimethylactyl chloride solution (0.03 ml, about 0.351 micro moles in dry CH₃CN/pyridine, v/v, 2 ml) were drawn into the syringe, which was agitated for five minutes (but for 15 minutes for cholesteryl H-phosphonate coupling). After each coupling, the reagents were ejected from the syringe and the procedure was continued by returning to the initial wash steps.

Oxidation, following final coupling step

and wash with dry CH₃CN (4.5 ml X 3), dry pyridine

(4.5 ml X 4) was performed with 0.1 58 in

CS₂/pyridine (1/1, v/v, 4.5 ml X 2), CH₃N (4.5 ml X

3), dry CH₃CN (4 ml X 3) and ether (5 ml X 4). After

drying the CPG-bound product was treated with 3.0 ml

concentrated NH₄OH at 55 degrees Centigrade for five

hours. Upon removal of NH₄OH by evaporation under

reduced pressure, CPG support was removed by

filtration; the filtrate freeze-dried overnight and

the product redissolved in 2.0 ml H₂O. For UV

spectroscopy, 10 micro liters of this solution were

added to 990 micro liters of H₂O.

HPLC data indicated about 50% of the reaction mixture was the desired product.

Spectroscopic methods also confirm the desired structure.

invention can also be utilized in a method for hybridizing with a complementary sequence in a solution under conditions conducive to the hybridization. Typically, these conditions are controlled by the complementary sequence. The specific conditions needed for hybridization would be known to one skilled in the art familiar with the complementary sequence and environment for hybridization.

The present invention further comprises the method of modifying the backbone of an oligonucleotide by the attachment of a fatty substance, preferably cholesteryl so that it will anchor into the cell membrane so that the modified oligonucleotide will hybridize with the complementary sequence. By anchoring into the cell membrane, the oligonucleotide may provide diagnostic or therapeutic activity. For example, the oligonucleotide compound 1 of Table 1 is complementary to the splice acceptor for site at 5349-5368 in HIV-1 and has been shown to inhibit replication of this virus in MoLT-3 cells by Zerial, et al., Nuc. Acids Res., 15:9909-9919 (1987) and Stein, et al., Cancer Res, 48:2659-2668 (1988).

variations of the basic sequence of compound 1.

Compounds 2 - 15 were designed to provide information on the antiviral properties of the cholesteryl modified oligonucleotides and, specifically, on the dependence of the antiviral activity on 1 (the number of cholesteryl fragments incorporated in the backbone chain); 2 (the nature of the main backbone section e.g. phosphodiester venus phosphorothicate links); 3 (the length of the oligonucleotide) and 4 (the sequence integrity of the oligonucleotide.

Samples of the oligomers were assayed in the following test:

Assays for HIV-1 Inhibition.

The inhibition of HIV-1 expression of H9 or

15 MOLT-3 cells in the presence of antisense

oligonucleotides was carried out by infecting 5 X 10⁵

cells per ml with 2.5-5 X 10⁸ virus particles of

HIV-1 strains HTLV-IIIB or HTLV-IIIC. Infection with

500-1000 virus particles per cell represents a

20 multiplicity of infection (MOI) of 0.5-1. HIV-1

infection of cells was carried out by simultaneous

addition of virus and cholesteryl modified oligomers

to the cells in culture. The cultures were incubated

in culture medium containing RPMI 1640, 10% (v/v)

25 fetal bovine serum, 2mM gultamine, and 250 micrograms

of gentamicin per ml, in a humidified atmosphere

containing 5% CO, at 37 degrees Centigrade. After four days, the cells and supernatant were examined for the level of HIV-1 expression by measuring syncytia (MOLT-3 cells) and viral antigen expression 5 as well as cell viability. The number of syncytia formed MOLT-3 cells were counted after triturating the cells to obtain an even distribution of the syncytia in the culture. The average number of syncytia as obtained by counting several fields in 10 duplicate cultures. Cell viability was measured in the presence of trypan blue, and the cells that excluded the dye were counted as viable cells. HIV-1 antigen expression was measured in cells fixed in methanol/acetone as described. Sarin, et al., 15 <u>Biochem. Pharmacol.</u>, <u>34</u>:075-4078 (1985) and Sarin, et al., J. Natl. Cancer Inst., 78:663-666 (1987). In brief, the cells were pelleted and then resuspended in phosphate-buffered saline (PBS) at a concentration of 106 cells per ml. The cells were spotted on 20 toxoplasmosis slides, air-dried, and fixed in methanol/acetone (1:1, v/v) for 15 minutes at room temperature. The slides were next incubated with 10% normal goat serum at room temperature for 30 minutes and washed with PBS four times. HIV-1 p24 or P17 25 monoclonal antibody was added to each well and the slides were incubated for 30 minutes in a humid

chamber 15 37 degrees Centigrade. The slides were then washed four times with PBS, incubated with fluorescein isothiocyanate-labeled goat anti-mouse IgG (Cappel Laboratories, Cochranville, Pennsylvania) 5 for 30 minutes at 37 degrees Centigrade, and then washed with PBS overnight. The slides were counterstained with Evan's blue, washed with PBS, mounted with 50% glycerol, and examined with a Zeiss fluorescence microsocpe. The percentages of cells 10 fluorescing in the oligomer-treated and untreated cultures were compared. Inhibition of HIV-1 expression in the presence of oligomers was found to be similar in both the H9 and the MOLT-3 cells.

Inhibition of HIV-1 expression and H9 and 15 MOLT-3 cells in the presence of cholesteryl modified oligonucleotides was carried out and results shown in Tables 2 and 3.

The data for the inhibition of formation of syncytia, an expression of HIV proteins P17, P24 and 20 reverse transcriptase shown for compounds 1 - 5 in Table 2 and compounds 6 - 16 in Table 3. The tables show results in ID₅₀ values for inhibition of syncytia (concentration of an oligomer in micro grams/ml. It gives 50% inhibition under the assay 25 condition) as an index.

The data from the tables describe favorable conclusions. The activity of the parent oligonucleotide, compound 1, is relatively low (ID50 less than 100). It appears that anchoring a 5 cholesteryl fragment to the oligonucleotide significantly enhances the anti-HTV activity (from ID50 greater than 100 to 10). Thus, the cholesteryl provides steroid means conjugated to the oligonucleotide for increasing the antiviral activity 10 of the oligonucleotide. Further, anchoring a second cholesteryl fragment does not appear to be an improvement because the second cholesteryl leads to a reduction in activity relative to the monocholesteryl-oligonucleotide. It appears that a 15 cholesteryl fragment to a phosphorothicate oligonucleotide analog enhances the antiviral property of the phosphorothicate derivative as shown in comparisons between compounds 6 and 7, 8 and 8, and 10 and 11. In the most favorable case, compound 20 7, the ID50 was reduced to 0.8 micrograms per milliter. With relatively large oligomers, those having 15 to 20-mers, the activity of the cholesteryl-oligonucleotides (natural phosphodiester links) appears to be independent of the chain link

(compare compounds 2 and 5). A lack of dependence of

activity on link has also been shown for unmodified

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oligonucleotides in the 15-20 mer range.

Additionally, the activity of the cholesteryl modified phosphorothicate derivates shows a downward trend as the length of the oligomer is decreased.

5 Thus, The ID50 values increase from 0.8 for the 20-mer (compound 7) to about 3.5 for the 10-15 mers (compound 11 and 9), to 13 for the 6-mer (compound 14).

From Tables 2 and 3, it can also be concluded that the anti-HIV activity of the 10 cholesteryl-modified oligonucleotides is not strongly dependent on the nucleotide sequence. conclusion applies both to the phosphodiester and the phosphorothicate cholesteryl derivatives (compared to 15 the data for compounds 2 and 3 has six mismatched base sites; and the data for compound 11 with that for compounds 12 and 13 which have 8 and 3 mismatched). For phosphorothicate derivatives, the activity of all three 10-mers is essentially the same 20 although the sequence is different. Further, the cholesteryl modified oligomers are not toxic to cells even at concentrations far in excess of those that lead to complete inhibition of the replication of HIV. For all derivates the LB50 was greater than 100 25 micrograms per ml.

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Additional Oligonucleotides having Steroids Conjugated Thereto Possessing Antiviral Activity

activity are set forth in Table 4. Activity is

5 reported as the average ID50 values (the concentration in micrograms per ug/ml of oligomer that leads to 50% inhibition of virus) determined by syncytia formation and expression of the viral proteins P17 and P24. The procedure is referenced in 10 Letsinger et al (1989) Proc. Natl. Acad. Sci. USA, 86:6553-6556. The abbreviations used in the Table are set forth below.

internucleoside phosphorothicate link, as in ... CsG...

over the company of the

25

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...N.N'... internucleoside link with a cholesteryl group tethered chol by a linker containing two carbon atoms, as in ..G.G..

...N.N'... internucleoside link with a cholesteryl group tethered by a linker containing ten carbon atoms, as in ..G.G..

CholesterylsN.. Terminal cholesteryl as in CholesterylsC...

CholesterylpC...

EXPERIMENTATION

All compounds were synthesized on controlled pore glass supports using a Biosearch 8600 5 DNA synthesizer or a syringe for manual manipulation. Internucleoside phosphorothioate links in compounds 1-4, 6, 8-16 were generated by conventional hydrogen phosphonate chemistry using 5'dimethoxytritylnucleosides and terminal oxidation of 10 all hydrogen phosphonate links with sulfur. Cholesteryl was joined to the chain in 1-4, 8, 10, 12, 14-16 by oxidative compling of 2-(cholesteryloxycarbonylamino)-ethylamine at an internucleoside hydrogen phosphonate link via 15 reaction with carbon tetrachloride. The procedure for tethering cholesteryl in compounds 11 and 13 was identical except that 2-(cholestery)oxycarbonylamino) -decylamine was used as the amine component. All these procedures are described in the 20 original patent application and/or reference 1. For synthesis of 6, cholesteryl was joined utilizing cholesteryl H-phosphonate. For the synthesis of compounds with phosphodiester links 5' to phosphorothicate links (compounds 5 and 7), chains 25 were built using methyl phosphoramidate coupling followed by oxidation with sulfur at each step. W. J.

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Stec, et al, Am. Chem. Soc., 106:6077 (1984). The terminal cholesteryl phosphate was added by oxidative coupling with cholesteryl H-phosphonate and carbon tetrachloride, as for compound 6, followed by standard oxidation with iodine/water. Treatment with thiophenol to remove the methyl protecting groups and ammonium hydroxide to remove base protecting groups and cleave the oligomer from the solid support the afforded the modified oligonucleotide.

- Compounds 15, 16 and a related substane,
 compound 17 with the structure XSXSXSXSXSXSXSXSXSXSX
 chol T,
 were prepared using the syringe technique and DMTOCH₂CH₂CH₂OP(O)(H)O (compound 18 in place of a
 nucleoside H-phosphonate. For synthesis of 18, 1,315 propanediol (75 mmol) was stirred with
- propanediol (75 mmol) was stirred with dimethoxytrityl chloride (15 mmol) in dry pyridine (25 ml) for about 20 hours at room temperature. Addition of water (100 ml), extraction with chloroform, and chromatography on a silica gel column
- 20 afforded 3-dimethoxyltrityloxypropanol as a viscous oil; 56% yield; NMR in CDCl₃: 1.85 (p, 2H, -CH₂-), 2.21 (broad t, 1H, 0-OH), 3.28 (broad t, 2H, -CH2(OH), 3.79 (broad t, 8H, two CH₃O-), 6.82 (q, 4H), 7.21-7.42 (m, 9H aromatic). This alcohol (6.9 mmol),
- 25 in dry acetonitrile (100 ml) was added dropwise over a period of 1 hour to phosphitilating solution

prepared from phosphorus trichloride (30 mmol), imidazol (98 mmol), and triethylamine (104 mmol) in acetonitrile (100 ml) at ice-bath temperature. After one hour of stirring the flask was allowed to warm to room temperature and to stand for two hours. Water (50 ml) was added and the mixture was concentrated under reduced pressure and extracted with chloroform. Chromatography on silica gel afforded 20.0 g (62%) of 18 as the triethylammonium salt; mp 132-135°C; NMR (CDCl₃): ppm 1.22 (t, 9H), 1.91 (p, 2H), 2.93 (q, 6H), 3.13 (t, 2H), 3.72 (S, 6H), 3.97 (t, 2H), 6.05 and 7.57 (singlets each, 1H), 6.77 (q, 4H), 7.14-7.39 (m, 9H).

Referring to Table 4, compound 1 is

complimentary to a region in HIV-I coating for a splice site for the TAT gene. Compound 3 is complimentary to a region for REV gene. Compound 4 is complimentary to a region for NEF gene.

Compounds 2, 5, 6, 7, 8 and 9 are all based 20 on sequence 1, with variations in position, mode of attachment, and the structure of the lipophilic steriod conjugate.

Compounds 10-14 are conjugates of homopolymers of nucleotides.

Referring to the activites shown in Table 4, compounds 1, 3, and 4, which are complimentary to various regions of the HIV-I virus, all show unusually high antiviral activity. These compounds may function by an antisense mechanism. Compound 3 is the most active cholesteryl-conjugated oligonucleotide that has been investigated.

Oligonucleotide phosphorothicate
derivatives bearing two cholesteryl groups, one
tethered at an internal position, are active, as
demonstrated by compounds 2 and 14. Activity for an
oligonucleotide with cholesteryl groups tethered at
the 5' and 3' terminal internucleoside positions as
set forth above it has been reported.

20 phosphodiester or a phosphorothicate link, are active. The activity is not strongly dependent on which of these links is used; that is, ID50 is 7 ug/ml for the phosphodiester, compound 7.

Linkage by the carbonylaminoethylamidate, such as in compound 8, affords a more active compound than linking by a phosphodiester, such as compound 7, or a phosphorothicate, such as compound 6.

Other highly lipophilic groups, such as stigmasteryl (compound 9) are comparable to cholesteryl in enhancing the antiviral activity of the oligonucleotide derivatives.

Cholesteryl conjugated homooligomers

10 containing thymidine, such as compounds 10, and 11,
or deoxycytidine, such as compounds 12, and 13, are
active antiviral pharmaceutical compositions. The
deoxycytidine oligomers are somewhat active than the
thymidine derivatives.

oligonucleotide having long carbon chains of for example 10 methylene groups, as well as short carbon chains of 2 methylene groups can be used in linking the cholesteryl to the phosphorus atom, as shown by comparing data for the thymidine derivatives (compounds 10 and 11) and the deoxycytidine derivatives (compounds 12 and 13. The compounds with the linker -NH(CH₂)₁₀NHCO- exhibit about the same activity as the corresponding compounds with the

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In view of the above, applicant has demonstrated the antiviral activity of the pharmaceutical composition of an oligonucleotide conjugated to a steriod. Applicant has shown that the oligonucleotide can be a homooligomer as well as a specific nucleotide sequence complimentary to various regions of a virus, such as the HIV-I virus. Applicant has also shown that various steriods conjugated to the oligomer have antiviral activity.

- The invention has been described in an illustrative manner, and it is to be understood that the terminology which has been used is intended to be in the nature of words of description rather than of limitation.
- Obviously, many modifications and variations of the present invention are possible in light of the above teachings. It is, therefore, to be understood that within the scope of the appended claims wherein reference numerals are merely for convenience and are not to be in any way limiting, the invention may be practiced otherwise than as specifically described.

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Table 2. Inhibition of HIV by Oligonucleotides with Cholesteryl Substituents

:ompound	conc.				
	μg/ml	syneytia	P24	Kľ	(Syncytla)
(control)	0.16	0			> 1.00
	0.8	· 3 ·			
	4	20			•
	20	34			
	100	45			
	2	0	0		10
	5	4	0	0	
	10	51	63	48	•
	20	95	88	-90	
	50	100	100	92	
	2	0	0	0	16
	5	2	13	0	
	10	22	70	0	
	20	77	69	٥	• •
•	50	100	100	84	
•	100	100	100	100	
	2	0	0	0	32
	5	3	0	0	
	10	7	0	0	
	20	28 -	32	26	
	50	85	88	75	
	100	100	100	100	
i	2	0	Q	0	11
	5	5	0	0	
	20	92	100	82	
	50	100	100	100	
	100	100	100	100	

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Table 3. Inhibition of HIV by Phosphorothicate Oligonucleotides with Cholesteryl Substituents

			•			
	μ g/ul :	syncycia	<u>8 inhibiri</u> P17	P24	RT	ID ₅₀ (syncytia)
6 (phospho-	2.5	15	13	22	33	6.0
diaster	6.25	56	67	81	70	•••
control)	10	90	89	89	85	
	25	100	100	100	100	
7	0.25	0	12	19	23	0.8
	1.0	74	69	70	68 .	••••
	1.5	100	100	100	100	
	6.0 _.	100	100	100	100	
8 (phosoho-	1.6	0	Ú	O	0	14.5
diester	6.25	15	16	26	26	.4.5
control)	25	95	84	82	67	
•	100	97	96	96 ·	72	
9	1.6	28	39	· 42	47	
	6.25	98	92	96		3_2
	25	98	96	96	73	
	100	98	96	11	76	
		70		96 .	88	
10 (phospho-	1.6	0	0	0	0	>100
diester	6.25	0	4	4	0	
control)	25	0	29	22	25	
	100	0	24	33	28	
ប	1.6	30	42	47	45	3.5
	6.25	97	86	88	61	
	25	97	92	92	70	
	100	98	92	96	89	
L 2	1.6	28	33	40		3.4
	6.25	93	60	67		3.4
	25	100	100	100		
	100	100	100	100		
L3	1.6	20	23	31		3.6
	6.25	89	56	67		J
	25	100	100	100		
	100	100	100	100		
.4	1.6	18	21	25		13
	6.25	35	30	32		·~
	25	90	70	66		
	100	100	100	100		
.5	25	80	20	25		
	50	99	99	90	•	

Table 4 New Modified Oligonucleotides

Cor	mpound St	ructure	ID50 (ug/ml)*
1•	AECEAECECE	Ceasaststscetegsasasasasteg_g	0.4-0.7
2	AsCsAsCsCs	CahahatatacatagahahahahatpG,G Chol chol	1.7
3	GeTeGeTeCs	Tacacacacatatacatatacacatagacaga	T 0.05
4	Asgstscscs	:Aetstsgsgstscststsasassgsgstsas	C.0 DakabaTaDaDaDaDaDaDaDaDaDaDaDaDaDaDaDaDaDaD
5	Cholestery	/lpascsascscsasaststscstsgsasas	
6	Cholestery	/lecstsgeasasasastsgsg	7
7	Cholestery	/lpcstsgsasasastsgsg	12
8	CSTSGSASA	BASASTSG _Z G Chol	1.5-2.5
9	Stigmaster	tylscetstsgsasasasatsgsg	7-14
10	TSTSTSTST:	STSTSTST_T Chol	10-18
11	. TsTsTsTsT	STSTSTST,T Žehol	15-30
1.2	? CsCsCsCsC	scscscsc c chol	· 6
13	CSCSCSCSC	scscscsc c Žchol	7
14	CSCSCSCSC	scsc_csc_c chol_chol	1.7 (toxic > 4 ug/ml)

a. A range in activity is shown for compounds tested more than once.
b. This compound, reported previously, is included for reference.

What is claimed is:

An oligonucleotide comprising:

wherein A is selected from the groups consisting of an aliphatic alkyl, branched aliphatic alkyl and a heteroatom containing an alkyl (branched) chain of 2 to 18 carbon atoms, R is selected from the group consisting of H and lower alkyl up to 12 carbon atoms; NUC is selected from the group consisting of an oligonucleotide and a phosphothicate oligonucleotide and B is a base.

20

2. An oligonucleotide as set forth in claim 1 wherein said oligonucleotide includes 7 to 20 nucleotides.

- A pharmaceutical composition consisting essentially of: an oligonucleotide and at least one steriod conjugated to said oligonucleotide for increasing the antiriral activity of said oligonucleotide.
 - 4. A composition as set forth in claim 3 wherein said steriod means is cholesterol conjugated to said oligonucleotide.

10

- 5. A composition as set forth in claim 4 wherein said oligonucleotides include phosphodiester bonds.
- 6. A composition as set forth in claim 5 including phosphothiate substitutions of phosphodiester bonds.
- 7. A composition as set forth in claim 3
 20 wherein said oligonucleotide is complementary to a nucleotide region of a virus.
 - 8. A composition as set forth in claim 7 wherein the virus is HIV-1 virus.

ŝ

- 9. A composition as set forth in claim 8 wherein said region of the HIV-1 virus is the TAT gene, the REV gene, or the NEF gene.
- 5 10. A composition as set forth in claim 3 wherein said steriod is stigmasteryl.
- 11. A composition as set forth in claim 3 including two steriods conjugated to said10 oligonucleotide.
 - 12. A composition as set forth in claim 3 wherein said oligonucleotide is a homopolymer.

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INTERNATIONAL SEARCH REPORT

International Application No PCT/US90/03204

I. CLASSIFICATION OF SUBJECT MATTER (if several classification symbols apply, indicate all) 2						
According to international Patent Classification (IPC) or to both National Classification and IPC						
INT CL. (5): A6IK 31/00; CO7H 15/00						
U.S. CL: 514/45,46,49.50; 536/27,29						
II. FIELDS SEARCHED						
Minimum Document Classification System i						
- Contraction of press.	lassification Symbols					
U.S. 514/45,46,49,50; 536/27,29						
Documentation Searched other the to the Extent that such Documents a	en Minimum Documentation are included in the Fields Searched s					
III. DOCUMENTS CONSIDERED TO BE RELEVANT 14						
Category *! Citation of Document, 16 with Indication, where appro	opmate, of the relevant passages 17 Relevant to Claim No. 18					
A,P Proceedings of the National Ace 86, September 1989, Robert I "Cholesterylconjugated oligonus properties, and activity as int of human immunodeficiency virus 6553-6556.	L. Letsinger et al., electides: Synthesis, mibitors of replication					
*A" document defining the general state of the art which is not considered to be of particular relevance "E" earlier document but published on or after the international filing date of priority date and not in conflict with the application but clied to understand the principle or theory underlying the invention invention which is cited to establish the publication date of another chainton or other special reason (as specified) "O" document referring to an oral disclosure, use, exhibition or other means "P" document published after the International filing date but later than the priority date claimed "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art. "A" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art. "A" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art. "A" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled to understand the principle or theory underlying the or priority date and not in conflict with the application but clied to understand the principle or theory underlying the invention cannot be considered to involve an inventive step when the document is combined with one or more other such document.						
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